

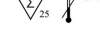
# Shigella Real Time PCR Kit User Manual For In Vitro Diagnostic Use Only







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### 1. Intended Use

Shigella real time PCR kit is used for the detection of Shigella in stool or water samples by using real time PCR systems

# 2. Principle of Real-Time PCR

The principle of the real-time detection is based on the fluorogenic 5'nuclease assay. During the PCR reaction, the DNA polymerase cleaves the probe at the 5' end and separates the reporter dye from the quencher dye only when the probe hybridizes to the target DNA. This cleavage results in the fluorescent signal generated by the cleaved reporter dye, which is monitored real-time by the PCR detection system. The PCR cycle at which an increase in the fluorescence signal is detected initially (Ct) is proportional to the amount of the specific PCR product. Monitoring the fluorescence intensities during Real Time allows the detection of the accumulating product without having to re-open the reaction tube after the amplification.

### 3. Product Description

Shigellosis is an infectious disease caused by a group of bacteria called Shigella. Most who are infected with Shigella develop diarrhea, fever, and stomach cramps starting a day or two after they are exposed to the bacteria. The diarrhea is often bloody. The Shigella germ is actually a family of bacteria that can cause diarrhea in humans. They are microscopic living creatures that pass from person to person. Shigella were discovered over 100 years ago by a Japanese scientist named Shiga, for whom they are named. Determining that Shigella is the cause of the illness depends on laboratory tests that identify Shigella in the stools of an infected person.

Shigella real time PCR kit contains a specific ready-to-use system for the detection of the Shigella (including S.boydii, S.dysenteriae, S.flexneri and S.sonnei) using PCR (polymerase chain reaction) in the real-time PCR system. The master contains reagents and enzyme for the specific amplification of the Shigella DNA. Fluorescence is emitted and measured by the real time systems' optical unit during the PCR. The detection of amplified Shigella DNA fragment is performed in fluorimeter **channel** 530nm with the fluorescent quencher BHQ1. In addition, the kit contains a system to identify possible PCR inhibition by measuring the 560nm fluorescence of the internal control (IC). An external positive control defined as  $1\times10^7$  copies/ml is supplied which allow the determination of the gene load. For further information, please refer to section 9.3 Quantitation.

### 4. Kit Contents

Ref.	Type of reagent	Presentation 25rxns	
1	DNA extraction buffer	2 vials, 1.5ml	
2	Shigella Reaction Mix	1 vial, 450μl	
3	PCR Enzyme Mix	1 vial, 12μl	
4	Molecular Grade Water	1 vial, 400µl	
5	Internal Control (IC)	1 vial, 30µl	
6	Shigella Positive control(1×10 <sup>7</sup> copies/ml)	1 vial, 30µl	

# Analysis sensitivity: 1×10<sup>3</sup> copies/ml;

LOQ:  $2 \times 10^3 \sim 1 \times 10^8$  copies/ml.

• Real time PCR system

· Refrigerator and Freezer

· Sterile filter tips for micro pipets

· Disposable gloves, powderless

· Vortex mixer

Cryo-container

Note: Analysis sensitivity depends on the sample volume, elution volume, nucleic acid extraction methods and other factors. If you use the DNA extraction buffer in the kit, the analysis sensitivity is the same as it declares. However, when the sample volume is dozens or even hundreds of times greater than elution volume by some concentrating method, it can be much higher.

# 5. Storage

- All reagents should be stored at -20°C. Storage at +4°C is not recommended.
- All reagents can be used until the expiration date indicated on the kit label.
- Repeated thawing and freezing (>3x) should be avoided, as this may reduce the sensitivity of the assay.
- · Cool all reagents during the working steps.
- Reaction mix should be stored in the dark.

# 6. Additionally Required Materials and Devices

- · Biological cabinet
- · Trypsin digestive Solution
- Real time PCR reaction tubes/plates • Pipets (0.5  $\mu l - 1000~\mu l)$
- Sterile microtubes
- · Biohazard waste container
- · Tube racks
- Desktop microcentrifuge for "eppendorf" type tubes (RCF max. 16,000 x g)

# 7. Warnings and Precaution

- · Carefully read this instruction before starting the procedure.
- · For in vitro diagnostic use only.
- This assay needs to be carried out by skilled personnel.
  Clinical samples should be regarded as potentially infectious materials and
- should be prepared in a laminar flow hood.
- This assay needs to be run according to Good Laboratory Practice.
- Do not use the kit after its expiration date.
- Avoid repeated thawing and freezing of the reagents, this may reduce the sensitivity of the test.
- Once the reagents have been thawed, vortex and centrifuge briefly the tubes before use.
  Quickly prepare the reaction mix on ice or in the cooling block.
- Set up two separate working areas: 1) Isolation of the RNA/ DNA and 2) Amplification/ detection of amplification products.

  • Pipets, vials and other working materials should not circulate among working units.
- Use always sterile pipette tips with filters.

# Wear separate coats and gloves in each area. Sample Collection, Storage and transportation

- Collect samples in sterile tubes;
- Specimens can be extracted immediately or frozen at -20°C to -80°C.
- Transportation of clinical specimens must comply with local regulations for the transport of etiologic agents

# 9. Procedure

# 9.1 DNA-Extraction

DNA extraction buffer is supplied in the kit. Please thaw the buffer thoroughly and spin down briefly in the centrifuge before use. It's better to use commercial kits for nucleic acid extraction

# 9.1.1 Stool samples

1) Take about 50mg samples to a 1.5ml tube; add 1.0ml normal saline then vortex vigorously.

- Centrifuge the tube at 13000rpm for 2 minutes, carefully remove and discard supernatant from the tube without disturbing the pellet.
- 2) Add 100µl DNA extraction buffer, close the tube then resuspend the pellet with vortex vigorously. Spin down briefly in a table centrifuge.
- 3) Incubate the tube for 10 minutes at 100°C.
- 4) Centrifuge the tube at 13000rpm for 5 minutes. The supernatant contains the DNA extracted and can be used for PCR template.

### 9.1.2 Water samples

- 1) Take 3 ml water to a tube, Centrifuge the tube at 13000rpm for 2 minutes, carefully remove and discard supernatant from the tube without disturbing the pellet.
- 2) Add 100µl DNA extraction buffer, close the tube then vortex for 10 seconds. Spin down briefly in a table centrifuge.
- 3) Incubate the tube for 10 minutes at 100°C.
- 4) Centrifuge the tube at 13000rpm for 5 minutes. The supernatant contains the DNA extracted and can be used for PCR template.

### Attention:

A. During the incubation, make sure the tube is not open, as the vapor will

volatilize into the air and may cause contamination if the sample is positive.

**B.** The extraction sample should be used in 3 hours or store at -20°C for one month.

C. Different DNA extraction kits are available. You may use your own extraction systems or the commercial kit based on the yield. For the DNA extraction, please comply with the manufacturer's instructions

#### 9.2 Internal Control

It is necessary to add internal control (IC) in the reaction mix. Internal Control (IC) allows the user to determine and control the possibility of PCR inhibition.

Add the internal control (IC) 1µl/rxn and the result will be shown in the 560nm.

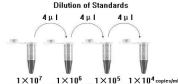
### 9.3 Quantitation

The kit can be used for quantitative or qualitative real-time PCR.

For performance of quantitative real-time PCR, Standard dilutions must prepare first as follows. Molecular Grade Water is used for dilution.

### The step of dilution is not needed for performance of qualitative real-time PCR.

Take positive control  $(1\times10^7\text{copies/ml})$  as the starting high standard in the first tube. Respectively pipette **36ul** of Molecular Grade Water into next three tubes. Do three dilutions as the following figures:



To generate a standard curve on the real-time system, all four dilution standards should be used and defined as standard with specification of the corresponding concentrations.

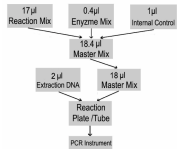
# Attention:

A. Mix thoroughly before next transfer.

B. The positive control contains high concentration of the target DNA. Therefore, be careful during the dilution in order to avoid contamination.

# 9.4 PCR Protocol

The Master Mix volume for each reaction should be pipetted as follows:



# \*PCR system without 560nm channel may be treated with 1μl Molecular Grade Water instead of 1μl IC.

- 1) The volumes of Reaction Mix and Enzyme Mix per reaction multiply with the number of samples, which includes the number of the controls, standards and sample prepared. Molecular Grade Water is used as the negative control. For reasons of unprecise pipetting, always add an extra virtual
- sample. Mix Master Mix completely then spin down briefly in a centrifuge.

  2) Pipet 18µl Master Mix with micropipets of sterile filter tips to each *Real time* PCR reaction plate/tube. Then separately add 2µl DNA sample, positive and negative controls to different reaction plate/tubes. Immediately close the plate/tubes to avoid contamination.
- 3) Spin down briefly in order to collect the Master Mix in the bottom of the reaction tubes.
- Perform the following protocol in the instrument:

37°C for 2min	1 cycle
94°C for 2min	1 cycle
93°C for 5sec, 60°C for 30sec	40cycles
(Fluorescence measured at 60°C)	40Cycles

Selecti	Selection of fluorescence channels		
530nm	Target Nucleic Acid		
560nm	IC		

10.Threshold setting: Choose Arithmetic as back ground and none as Noise Band method, then adjust the Noise band just above the maximum level of molecular grade water, and adjust the threshold just under the minimum of the positive control.

11.Calibration for quantitative detection: Input each concentration of standard controls at the end of run, and a standard curve will be automatically formed.

12.Quality control: Negative control, positive control, internal control and QS curve must be performed correctly, otherwise the sample results is invalid.

Channel	Crossing point value				
Control	530nm	560nm			
Molecular Grade Water	Blank	25~35			
Positive Control(qualitative assay)	≤35				
QS (quantitative detection)	Correlation coefficien	t of QS curve≤-0.98			

# 13. Data Analysis and Interpretation

le ionowing results are possible.					
	Crossing point value		Result Analysis		
	530nm	560nm	Result Allarysis		
1#	Blank	25~35	Below the detection limit or negative		
2#	≤35		Positive; and the software displays the quantitative value		
3#	35~40	25~35	Re-test; If it is still 35~40, report as 1#		
4#	Blank	Blank	PCR Inhibition; No diagnosis can be concluded.		
	2#	530nm 1# Blank 2# ≤35 3# 35~40	530nm 560nm 1# Blank 25~35 2# ≤35 — 3# 35~40 25~35		

For further questions or problems, please contact our technical support at trade@liferiver.com.cn